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Substitute for form 1449/PTO				Complete If Known		
00.	5511.01.01117 144671 10			Application Number	10/539,180-Conf. #2257	
11	NFORMATION	N DI	SCLOSURE	Filing Date	February 28, 2005	
S	TATEMENT I	BY A	APPLICANT	First Named Inventor	Blas Cerda	
				Art Unit	1657	
	(Use as many sh	eets a	s necessary)	Examiner Name	P. C. Martin	
Sheet	Sheet 1 of 2		Attorney Docket Number	NEN-23002/16		

U.S. PATENT DOCUMENTS							
Examiner Initials*	Cite No.1	Document Number Number-Kind Code ² (<i>if known</i>)	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages or Relevar Figures Appear		
	AA*	US-5,096,812	03-17-1992	Rachel et al			
	AB* US-6,258,605		07-10-2001	Chace			
	AC*	US-6,455,321	09-24-2002	Chace			
	AD*	US-6,670,194	12-30-2003	Aebersold et al			
	AE*	US-5,629,210	05-13-1997	Hercules et al			
	AF*	US-5,719,035	02-17-1998	Rosenthal et al			

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Examiner Initials*	Cite No.1	Foreign Patent Document Country Code ³ -Number ⁴ -Kind Code ⁵ (# known)	Publication Date MM-DD-YYYY	Name of Patentee or Applicant of Cited Document	Pages, Columns, Lines, Where Relevant Passages Or Relevant Figures Appear			

		NON PATENT LITERATURE DOCUMENTS					
Examiner Initials	Cite No.1	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published.	T²				
	CA	Online Mendelian Inheritance in Man (OMIM) entry for Zellweger syndrome. 2005. Accessed1 11/29/06; 13 pages					
	СВ	OMIM entry for Kears-Sayre syndrome. 2006. Access online 11/29/06. 13 pages					
	СС	J.O. Sass et al., "Mutations in ACY1, the Gene Encoding Aminoacylase 1, Cause a Novel Inborn Error of Metabolism," Am. J. Hum. Genet. 2006, 78:401-409					
	CD	K.L. Peterson, D.K. Srivastava; "Functional role of a distal (3'-phosphate) group of CoA in the recombinant human liver medium-chain acyl-CoA dehydrogenase-catalysed reaction," Biochem. J 325: 751-60					
	CE	R. Ramsay et al., "Carnitine palmitoyltransferase in human erythrocyte membrane," Biochem. J., 1991, Vol. 275, pp. 685-88					
	CF	Definition of "metabolism," 1991. Webster's College Dictionary, Random House, page 851					
	CG	P. Rinaldo et al.; "Disorders of fatty acid transport and mitochondrial oxidation: Challenges and dilemmas of metabolic evaluation," Nov/Dec 200, Vol. 2, No. 6, pp. 338-44					
	СН	P. Rinaldo et al., "Fatty Acid Oxidation Disorders," Annu. Rev. Physiol., 2002, 64: 477-501					
	CI	M. McCaman et al., "Fluorimetric Method for the Determination of Phenylalanine in Serum," J. Lab. Clin. Med., Vol. 59, No. 5, Aug. 1961, pp. 885, 887, 889.					
	CJ	N. Chamoles et al., "Hurler-like Phenotype: Enzymatic Diagnosis in Dried Blood Spots on Filter Paper," Clinical Chemistry, 47:12, pp. 2098-2102					

Examiner		Date	
Signature		Considered	

^{*}EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant.

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ск	N. Chamoles et al., "Fabry Disease: Enzymatic Diagnosis in Dried Blood Spots on Filter Paper, Clinica Chemica Acta 308, 2001, p. 195-96	
CL	L. Sweetman, "Newborn Screening by Tandem Mass Spectrometry (MS-MS), Clinical Chemistry, Vol. 47, No. 11, 2001, pp. 1937-38	
СМ	A. Fujimoto et al., "Quantitative Beutler Test for Newborn Mass Screening of Galactosemia Using a Fluorometric Microplate Reader, Clinical Chemistry 46:6, 2000, pp. 806-10	
CN	D. Chase, "Rapid diagnosis of homocystinuria and other hypermethioninemias from newborns' blood spots by tandem mass spectrometry," Clinical Chemistry, 42:3, 1996, pp. 349-55	
CO	Fisher Technical Assistance website: solvent selection guide	
СР	B. Im et al., "Bacterial Degradation of Biotin," Vol. 248, No. 22, Nov. 1973, pp. 7798-805	
	CK CL CM CN CO	the item (book, magazine, journal, serial, symposium, catalog, etc.), date, page(s), volume-issue number(s), publisher, city and/or country where published. CK N. Chamoles et al., "Fabry Disease: Enzymatic Diagnosis in Dried Blood Spots on Filter Paper, Clinica Chemica Acta 308, 2001, p. 195-96 CL L. Sweetman, "Newborn Screening by Tandem Mass Spectrometry (MS-MS), Clinical Chemistry, Vol. 47, No. 11, 2001, pp. 1937-38 CM A. Fujimoto et al., "Quantitative Beutler Test for Newborn Mass Screening of Galactosemia Using a Fluorometric Microplate Reader, Clinical Chemistry 46:6, 2000, pp. 806-10 CN D. Chase, "Rapid diagnosis of homocystinuria and other hypermethioninemias from newborns' blood spots by tandem mass spectrometry," Clinical Chemistry, 42:3, 1996, pp. 349-55 CO Fisher Technical Assistance website: solvent selection guide

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